ID NO: 6 INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN-LIKE POLYPEPTIDE (IDENTIFIED AS IGFBP-7-HY1) WITH MUS MUSCULUS INSULIN-LIKE GROWTH BLASTP ALIGNMENT OF SEQ

FACTOR BINDING PROTEIN SEQ IN NO:13 Query: IGFBP_like protein (SEQ ID NO: 6)

ID NO: SEQ >gi | 9055246 (AB006141) IGFBP-like protein [Mus musculus] Subject:

13

Length = 27

Score = 1170 (416.9 bits), Expect = 1.8e-118, P = 1.8e-118 209/269 (77%), Positives = 232/269 (86%) ll Identities

67 LPLLLLLLLLPPLSPSLGIRDVGGRRPKCGPCRPEGCPAPAPCPAPGISALDECGCCA ω Query:

+P L LLLL LLP L+ LG+RD G R P+C PC+ + CPAP+PCPAP ISA DECGCCA

50 MPRLPLLLL-LLPSLARGLGLRDAGRRHPECSPCQQDRCPAPSPCPAPWISARDECGCCA Sbjct:

127 RCLGAEGASCGGRAGGRCGPGLVCASQAAGAAPEGTGLCVCAQRGTVCGSDGRSYPSVCA 68 Query:

119 RCLGAEGASCGGPVGSRCGPGLVCASRASGTAPEGTGLCVCAQRGAVCGSDGRSYSSİCA RCLGAEGASCGG G RCGPGLVCAS+A+G APEGTGLCVCAQRG VCGSDGRSY S+CA 9

Sbjct:

187 LRLRARHTPRAHPGHLHKARDGPCEFAPVVVVPPRSVHNVTGAQVGLSCEVRAVPTPVIT 128 Query:

179 LRIRARHAPRAHHGHLHKARDGPCEFAPVVLMPPRDIHNVTGTQVFLSCEVKAVPTPVIT LRLRARH PRAH GHLHKARDGPCEFAPVV++PPR +HNVTG QV LSCEV+AVPTPVIT 120 Sbjct:

247 WRKVTKSPEGTQALEELPGDHVNIAVQVRGGPSDHEATAWILINPLRKEDEGVYQCHAAN 188 Query:

SPEGT+ LEELPGDHVNIAVQVRGGPSDHE T+WILINPLRKEDEGVY CHAAN W+KV

239 WKKVKHSPEGTEGLEELPGDHVNIAVQVRGGPSDHETTSWILINPLRKEDEGVYHCHAAN 180 Sbjct:

Query: 248 MVGEAESHSTVTVLDLSKYRSFHFPAPDD 276

+GEA+SH TVTVLDL++Y+S + P D

Sbjct: 240 AIGEAQSHGTVTVLDLNRYKSLYSSVPGD 268

BLASTP ALIGNMENT OF SEQ ID NO: 6 INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN-LIKE POLYPEPTIDE (IDENTIFIED AS IGFBP-7-HY1) WITH HOMO SAPIENS PROSTOGLANDIN I2 NO:14

IGFBP like protein (SEQ ID NO: 6) Query:

14) >gi|1082724 Prostoglandin I2 [Homo sapiens (SEQ ID NO: Subject:

Length:

7.9e-61 Identities = 123/273 (45%), Positives = 150/273 (54%) Score = 570 (205.7 bits), Expect = 7.9e-61, P =

54 PSLRALLLGAAGLLLLLL - PLSSSSS-SDT - - - - - CGPCEPASCPPLPPLGCLLGETR 4 Query:

CGPC P CP LLLL L PLS S P L +LLL

9 PRL-SLLLPLLLLLLLPLLPPLSPSLGIRDVGGRRPKCGPCRPEGCPAPAPCPAPG1SAL a Sbjct DACGCCPMCARGEGEPCGGGGAGRGYCAPGMECVKSRKRKGKAGAAAGGPGVSGVCVCK 55 Query:

+G+QAC AGAA G D CGCC C BG CGG GR C PG+ C

109 DECGCCARCLGAEGASCGGRAGGR--CGPGLVCASQA-----AGAAPEG---TGLCVCA 61 Sbjct:

174 SRYPVCGSDGTTYPSGCQLRAASQRAESRGEKAITQVSKGTCEQGPSIVTPPKDIWNVTG 115 Query:

P +V PP+ + NVTG G CE R VCGSDG +YPS C LR ++

169 110 ORGTVCGSDGRSYPSVCALRLRARHTPRAHPGHLHKARDGPCEFAPVVVVPPRSVHNVTG Sbjct:

234 175 AQVYLSCEVIGIPTPVLIWNKVKRGHYGVQRTELLPGDRDNLAIQTRGGPEKHEVTGWVL Query:

AQV LSCEV +PTPV+ W KV + G Q E LPGD N+A+Q RGGP HE T W+L

170 AQVGLSCEVRAVPTPVITWRKVTKSPEGTQALEELPGDHVNIAVQVRGGPSDHEATAWIL 229 Sbjct:

235 VSPLSKEDAGEYECHASNSQGQASASAKITVVD 267 Query:

++PL KED G Y+CHA+N G+A + + +TV+D

262 INPLRKEDEGVYQCHAANMVGEAESHSTVTVLD 230 Sbjct: